

# BEST AVAILABLE COPY

		tG	CI	'AT	CTC	cc	TT	rGCr	CAG	GGC	GC	AG	AGA	AA7	rga'	TG	gaç	GT	GAA	TAG	CAC	.17	rgc	TIC	GA	60
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	61	AA	CG	rcc	TG	GC	AC	CAT	GGI	TAT	GC.	MG	TCI	cc	TCC	TA.	.CG	TGI	CCI	YETY	CGC	GT.	AGA	GT	CCC	
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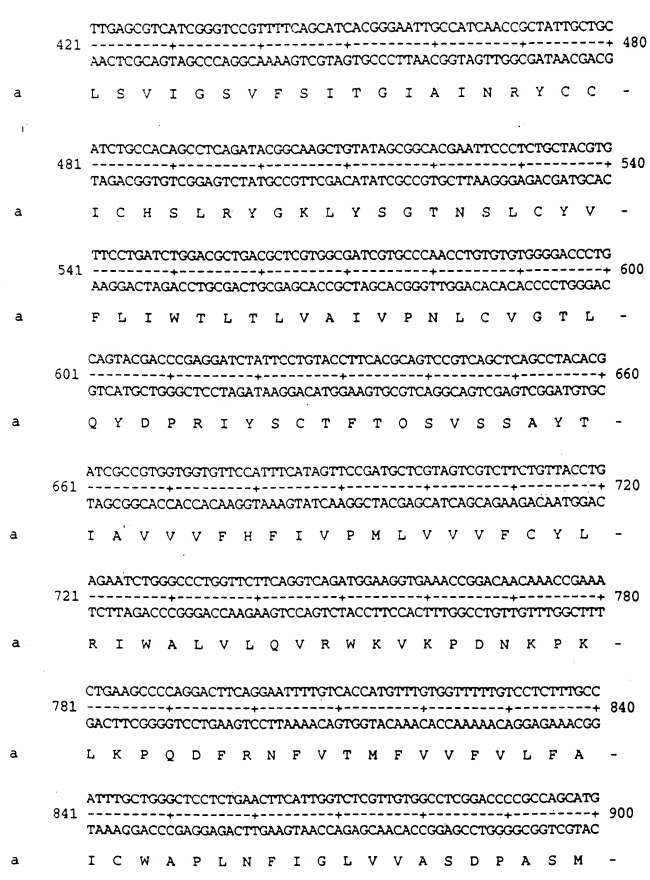
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	721	CA	\GC	\AA	\GT	rga(	CACA	\AA(	CAGA	CT	rga(	SAAZ	\TT	rcm	rga	CCA	TGT	TIC	TC	TC:	MIG	T
	, 21	GITI	CGI	(ALA)	ICA/	ACTO	गुज	TTC	TCI	'GA/	CTC	TT	'AA	\GA/	CT	GGT	ACA	AAC	ACC	CAG	AAAC	+ 780 A
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	901		TAA 	CAG	TTG	TCT	CAA			TAT.					AAA	TCA	AAA	CT	rcc		AGGA	960
		AAA	ATT	GTC	AAC	AGA	GTT.								TT	AGT	TTT	'GA/	AGGC		CCT	
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	961	CAT			+			-+-			+				+			-+-			+	1020
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	1021	TCC	TCC	TTG	+ ACT	ccc	TAA	CTT	TTC	ATT	:GG/	AGC	GG	rcg/	ACA	TTG	GTT	-+- GTT	GIT	AGT		1080
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	AGATATICTAGGAGAAGGTCACTIGTGGCTGAGCAGGAAATGGTGCGAAAATGGT 1081+ 1140 TCTATACGATCCTCTTCGTTCCAGTGACACCGACTCTTTTACCACGCTTTTACCA	D M L G E A R S L W L S R R N G A K M V -	GATCATCAGGCCAAGAAAAGCACAAATTGCAATCATCCATC	IIRPRKAQIAIIHQIFWPQ -	GAGTTCATGGGCAACATGCCGTCAAGACTTACCGGAGAAGATGGCTGCCG 1201+ 1260 CTCAAGTACCCGTTGTACGGCAGTTCTGTGTTTCTAATGGCCTCTTCTTCTACCGACGGC	SSWATCRODTKITGEEDGCR-	TGAACTGTGCAAGGAGTTTCCCAAAGGTGAGCCCAATGCACTATATCCACATTAT SEQ ID NO: 1 1261+ 1320 ACTTGACACGTTCCTGCCCTAAAGGGTTTCCACTCTGGGTTACGTGATATAGGTGTAATA	ELCKDGISQR * SEQIDNO:2 FIG. 1d	
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241				-+-			+			GCT CGA	+			+				+			+	300
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361				-+-			+			CCT	+			-+-				+			+	420
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FIG. 2a



901		ACC	CAG	GAT	CCC	CCGA	AGTY	GGC'					GTT	ACT	ATA:	rgg	CAT	ATT	TCA	ACAG	C + 960
901		TGG	GTC	CTA	GGC	GC1	CAC	CG					CAA	rga'	rat?	ACC	GTA'	TAA	AGT	TGTC	
	A	P	R	I	P	E	W	L	F	V	A	S	Y	Y	M	Ά	Y	F	N	S	-
961				-+-			+				+			-+-			+				- 1020
÷	AC	GGA	GTT.	ACG	CTA	GTA	TAT	ACC	TGA	ATG	CTI	GGI	LLLL	'AAA'	GTC	CG1	CCI	TAT	GT	CIALAI	ŗ
	C	L	N	A	I	I	Y	G	L	L	N	Q	N	F	R	Q	E	, <b>Y</b>	R	K	-
1021		TATA	AGTC	TC	ATT	GTG'	TAC	CAC	CAA	GAT	GTT	CTT	rgr	GGA'	rag(	CTC	CAA	rca	TGT.	AGCA	1080
1021		TAT	CAC	JAG'	raa	CAC	ATG	GTG	GTT	CTA	CAA	GAA	ACA(	CCT	ATC(	3AG(	STT	AGT.	ACA'	TCGT	
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	CT	ATC:	TAZ	\TT	rgc	GTT	TGG	GAG	AGG	TAA'	TTA'	rcg(	GTT	GGT?	ATTO	GA?	TA?	LaTaIA	CCA	CCTG	
	D	R	I	K	R	K	P	s	P	L	I	A	N	Н	N	L	I	K	V	D	-
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	961	901 CG A  7G 961 AC C  L021 TA  I  L081 CT D	901 CGTGG  A P  TGCCT  961 ACGGA  C L  ATTATA TAATATA I I  1021 TAATATA CTATCT D R  TCCGTT	GATAGAATT  GATACTTAA  GATACTTAA  CTATCTTAA  TCCGTTTAA	GATAGAATTAA  GATACTTAATT  D R I  CGTGGGTCCTA  A P R I  TGCCTCAATGO  TGCGTCAATGO  C L N A  ATTATAGTCTCA  TAATATCAGAG  I I V S  CTATCTTAATT  D R I K  TCCGTTTAA SE  141 11	CGTGGGTCCTAGGC  A P R I P  TGCCTCAATGCGAT  961 ACGGAGTTACGCTA  C L N A I  ATTATAGTCTCATT  TAATATCAGAGTAA  I I V S L  GATAGAATTAAACG  CTATCTTAATTTGC  D R I K R  TCCGTTTAA SEQ I  141	CGTGGGTCCTAGGGGCT  A P R I P E  GGTGGGTCCTAGGGGCT  A P R I P E  GGTGGGTCCTAGGGGCT  TGCCTCAATGCGATCAT  C L N A I I  ATTATAGTCTCATTGTGT  TAATATCAGAGTAACAC  I I V S L C  GATAGAATTAAACGCAA  CTATCTTAATTTGCGTT  D R I K R K  TCCGTTTAA SEQ ID N  141 1149	CGTGGGTCCTAGGGGCTCAG  A P R I P E W  TGCCTCAATGCGATCATATA  961	CGTGGGTCCTAGGGGCTCACCG  A P R I P E W L  GCCTCAATGCGATCATATATGC  ACGGAGTTACGCTAGTATATACC  C L N A I I Y G  ATTATAGTCTCATTGTGTACCAC  TAATATCAGAGTAACACATGGTG  I I V S L C T T  GATAGAATTAAACGCAAACCCTC  CTATCTTAATTTGCGTTTGGGAG  D R I K R K P S  TCCGTTTAA SEQ ID NO:3	CGTGGGTCCTAGGGGCTCACCGACA  A P R I P E W L F  GCCTCAATGCGATCATATATGGACT  GC L N A I I Y G L  ATTATAGTCTCATTGTGTACCACCAA  ATTATACAGAGTAACACATGGTGGTT  I I V S L C T T K  GATAGAATTAAACGCAAACCCTCTCC  CTATCTTAATTTCCGTTTGGGAGAGG  D R I K R K P S P  TCCGTTTAA SEQ ID NO:3	CGTGGGTCCTAGGGGCTCACCGACAAACA  A P R I P E W L F V  GCCTCAATGCGATCATATATGGACTACT  961 ACGGAGTTACGCTAGTATATACCTGATGAT  C L N A I I Y G L L  ATTATAGTCTCATTGTGTACCACCAAGATA  TAATATCAGAGTAACACATGGTGGTTCTAA  I I V S L C T T K M  GATAGAATTAAACGCAAACCCTCTCCATTA  CTATCTTAATTTCCGTTTGGGAGAGGTAAA  D R I K R K P S P L  TCCGTTTAA SEQ ID NO:3  141	CGTGGGTCCTAGGGGCTCACCGACAAACACCC  A P R I P E W L F V A  1021  ACGGAGTTACGCTAGTATATAGGACTACTGAA  ACGGAGTTACGCTAGTATATACCTGATGACTT  C L N A I I Y G L L N  ATTATAGTCTCATTGTGTACCACCAAGATGTT  TAATATCAGAGTAACACATGGTGGTTCTACAAC  I I V S L C T T K M F  GATAGAATTAAACGCAAACCCTCTCCATTAATA  CTATCTTAATTTGCGTTTGGGAGAGGTAATTAC  D R I K R K P S P L I  TCCGTTTAA SEQ ID NO:3	CGTGGGTCCTAGGGGCTCACCGACAAACACCGATC  A P R I P E W L F V A S  TGCCTCAATGCGATCATATATGGACTACTGAACCA  961 ACGGAGTTACGCTAGTATATACCTGATGACTTGGT  C L N A I I Y G L L N Q  ATTATAGTCTCATTGTGTACCACCAAGATGTTCTT  TAATATCAGAGTAACACATGGTGGTTCTACAAGAA  I I V S L C T T K M F F  GATAGAATTAAACGCAAACCCTCTCCATTAATAGCC  D R I K R K P S P L I A  TCCGTTTAA SEQ ID NO:3  141	CGTGGGTCCTAGGGGCTCACCGACAAACACCGATCAA'  A P R I P E W L F V A S Y  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAA  961	CGTGGGTCCTAGGGGCTCACCGACAAACACCCGATCAATGA'  A P R I P E W L F V A S Y Y  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAATTT  961 ACGGAGTTACGCTAGTATATACCTGATGACTTGGTTTTAAA  C L N A I I Y G L L N Q N F  ATTATAGTCTCATTGTGTACCACCAAGATGTTCTTTGTGGA'  TAATATCAGAGTAACACATGGTGGTTCTACAAGAAACACCTT  I I V S L C T T K M F F V D  GATAGAATTAAACGCAAACCCTCTCCATTAATAGCCAACCACCACCACCACCACCACCACCACCACCACCAC	CGTGGGTCCTAGGGGCTCACCGACAAACACCGATCAATGATATA  A P R I P E W L F V A S Y Y M  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAATTTCAG  961 ACGGAGTTACGCTAGTATATACCTGATGACTTGGTTTTAAAGTC  C L N A I I Y G L L N Q N F R  ATTATAGTCTCATTGTGTACCACCAAGATGTTCTTTGTGGATAGC  I I V S L C T T K M F F V D S  GATAGAATTAAACGCAAAACCCTCTCCATTAATAGCCAACCATAAC  1081 CTATCTTAATTTGCGTTTGGGAGAGGTAATTATCGGTTGGTATTC  D R I K R K P S P L I A N H N  TCCGTTTAA SEQ ID NO:3  .141	CGTGGGTCCTAGGGGCTCACCGACAAACACCGATCAATGATATACCC  A P R I P E W L F V A S Y Y M A  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAATTTCAGGCA  ACGGAGTTACGCTAGTATATACCTGATGACTTGGTTTTAAAGTCCGT  C L N A I I Y G L L N Q N F R Q  ATTATAGTCTCATTGTGTACCACCAAGATGTTCTTTGTGGATAGCTCC  TAATATCAGAGTAACACATGGTGGTTCTACAAGAAACACCTATCGAGC  I I V S L C T T K M F F V D S S  GATAGAATTAAACGCAAAACCCTCTCCATTAATAGCCAACCATAACCTA  CTATCTTAATTTGCGTTTGGGAGAGGTAATTATCGGTTGGTATTGGAT  D R I K R K P S P L I A N H N L  TCCGTTTAA SEQ ID NO:3  111 1149	901 CGTGGGTCCTAGGGGCTCACCGACAAACACCGATCAATGATATACCGTA' A P R I P E W L F V A S Y Y M A Y  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAATTTCAGGCAGG	901 CGTGGGTCCTAGGGGCTCACCGACAAACACCGATCAATGATATACCGTATAA  A P R I P E W L F V A S Y Y M A Y F  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAATTTCAGGCAGG	CGTGGGTCCTAGGGGCTCACCGACAAACACCGATCAATGATATACCGTATAAAGT  A P R I P E W L F V A S Y Y M A Y F N  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAATTTCAGGCAGG	CGTGGGTCCTAGGGGCTCACCGACAAACACCGATCAATGATATACCGTATAAAGTTGTC  A P R I P E W L F V A S Y Y M A Y F N S  TGCCTCAATGCGATCATATATGGACTACTGAACCAAAATTTCAGGCAGG

FIG. 2c

1	ΑT	GAA	GGG	CAA	TGT	CAG	CGA	\GC'I			ATC				\GG(	CTC	CAG	GCG	GCG	GGGAG	60
•	TA	CTT)	ccc	GTT	ACA	GTC	GCI	CGA							CCC	GAG	GTC	CGC	CGC	CCCTC	
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61	GG 	AGC	GAG	ACCI	ACG.	ACC	GTC	CTC	GAT	GCC	CTC	TAC	ACT	GGC	CT	CA	rcc	rca'	TCT	ITACC AATGG	120
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121																		<b>_</b>		AGAAG + ICTTC	180
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181				- · <del>-</del>			+				+			-+-			+			GGCT + CCGA	240
a	L	R	N	S	G	N	I	F	V	v	s	L L	A	v	A	D	L.	٧	٧	A	<b>-</b>
241											<b>+</b>			-+-						ATAT	300
241	CA	AAT	GGG	ATA	AGG(	CAA	CCA	CGA	CTG	TAG	АТА	GGA	PTA	GTT	GCC	TAC	CTI	'AGA	'CCC	TATA	
a	V	Υ,	P	Y	P	L	٧	L	Т	S	I	L	N	И	G	W	N	L	G	Y	-
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361				-4-			+				+			- 1			+			CAAA	420
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A21		ATA	CAG	TAA	CAA	GAA	CTC	GCT	CTG	СТА	CGT	GTT	CCI	CAT	ATG	GAT	GCI	GAC	ACT	CATC	480
423				-+-		·					+			-+-			+			CATC  GTAG	430

481	GC	CAT	CAT	GCC(	CAA	CCT	GCA +	AAC	CGG	AAC	ACT	CCA	GTA	ACGA	TCC	CCC	GA'	CT	CTC	CTGT	540
	CG	GTA	GTAC	CGG	GTT(	CGA	CGI	TTC	GCC	TTC	TGA	lgg1	CAT	rgci	CAGC	CCC	CT	AGA	TGA	GGACA	
	A	I	М	P	N	L	Q	τ	G	T	L	Q	Y	D	₽	R	I	Y	<b>.</b> \$	С	-
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991	TCT	rgcc	TCC	CAC	TT	rcc	GCT(	GII	GTT	CGG	+ GTT	TGA	CTIV	-+- CGG	ggt(	CCT	GAA	GTC	CTT	GAAA	720
	R	R	R	V	K	P	D	N	K	P	K	L	K	P	Q	D	F	R	N	F	-
721				+	'-		+-				+			-+-			+			+	780
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	GG!	rct:	rat?	rgte	GCC	etc <i>i</i>	AGA(	ا د	TGC	CAC	САТ	GGT	ccc	CAG	Gat	ccc	AGA	<b>IGT</b> C	GCT	'GTTC	
781							+				+			-+-			+	. – – -		+	840
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	541 601 721 781	481 CG  A  541 TGC  T  601 CAC  V  781 CAC  G  841 CAC	GGGTAG  A I  ACCTTO TGGAAG  T F  GTGCCT CACGGA  V P  AGACGC R R  721 CAGTGC V T  781 CCAGAG G L  841 CACCGG	CGGTAGTAGA  A I M  ACCTTCACG TGGAAGTGG T F T  GTGCCTATG CACGGATAG V P M  AGACGGAGG TCTGCCTCC R R R  721 CAGTGGTAG V T M  781 GGTCTTATT CCAGAATAG G L I  841 GTGGCTAGT	CGGTAGTACGCA  A I M P  ACCTTCACCCAC  TGGAAGTGGGTC  T F T Q  GTGCCTATGATT  CACGGATACTAA  V P M I  AGACGGAGGGTC  R R R V  721  CAGTGGTACAAC  V T M F  CCAGAATAACAC  G L I V  841  CACCGATCAACCA	CGGTAGTACGGGTT  A I M P N  ACCTTCACCCAGTC  TGGAAGTGCGTCAG  T F T Q S  GTGCCTATGATTATT  CACGGATACTAATAA  V P M I I  AGACGGAGGGTGAA  AGACGGAGGGTGAA  OF TCTGCCTCCCACTT  R R R V K  721  CAGTGGTACAAGCA  V T M F V  781  GGTCTTATTGTCGCC  G L I V A  GTGGCTAGTTACTAC  841  CACCGATCAATGATC  CACCGATCAATGATC  841  CACCGATCAATGATC  CACCGATCAATGATC  CACCGATCAATGATC  CACCGATCAATGATC	CGGTAGTACGGGTTGGA  A I M P N L  ACCTTCACCCAGTCTGT TGGAAGTGGGTCAGACA T F T Q S V  GTGCCTATGATTATTGTC CACGGATACTAATAACAC V P M I I V  AGACGGAGGGTGAAACCC R R R V K P  GTCACCATGTTCGTAGT CAGTGGTACAAGCATCA V T M F V V  GGTCTTATTGTCGCCTC GGTCTTATTGTCGCCTC CAGAATAACACCCGGAG G L I V A S  GTGGCTAGTTACTACCTC S41  CACCGATCAATGATGATGGAC  GTGGCTAGTTACTACCTC CACCGATCAATGATGGAC  GTGGCTAGTTACTACCTC CACCGATCAATGATGGAC	CGGTAGTACGGGTTCGACGI  A I M P N L Q  ACCTTCACCCAGICTGTCAG  TGGAAGTGGGTCAGACAGTC  T F T Q S V S  GTGCCTATGATTATTGTCATA  CACGGATACTAATAACAGTAA  V P M I I V I  AGACGGAGGGTGAAACCCGAA  TCTGCCTCCCACTTTGCGCTA  R R R V K P D  721  GTCACCATGTTCGTAGTTIT  CAGTGGTACAAGCATCAAAA  V T M F V V F  GGTCTTATTGTCGCCTCAGAA  781  GGTCTTATTGTCGCCTCAGAA  CCAGAATAACACCGGAGTCTA  G L I V A S D  GTGGCTAGTTACTACCTGGCGAAACCCGAAACACCCGGAGTCTAAAACACCCGGAGTCTAAAACACCGGAGTCTAACACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACACCGGAGTCTAAAAACACCGGAGTCTAAAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACACGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACACCGGAGTCTAAAACAACAACAACAACAACAACAACAACAACAACAA	CGGTAGTACGGGTTCGACGTTC  A I M P N L Q T  ACCTTCACCCAGTCTGTCAGCTC  TGGAAGTGGGTCAGACAGTCGAG  T F T Q S V S S  GTGCCTATGATTATTGTCATCTT  CACGGATACTAATAACAGTAGAA  V P M I I V I F  AGACGGAGGGTGAAACCCGACAA  AGACGGAGGGTGAAACCCGACAA  R R R V K P D N  GTCACCATGTTCGTAGTTTTTGTC  CAGTGGTACAAGCATCAAAAACA  V T M F V V F V  GGTCTTATTGTCGCCTCAGACCC  G L I V A S D P  GTGGCTAGTTCATACTACCTGCGGTAG  841  CACCGATCAATGATGGACCGCATA	CGGTAGTACGGGTTCGACGTTGGCC  A I M P N L Q T G  ACCTTCACCCAGTCTGTCAGCTCAGCTCAGCTCAGCT	CGGTAGTACGGGTTGGACGTTGGCCTTC  A I M P N L Q T G T  ACCTTCACCCAGTCTGTCAGCTCAGCGTA TGGAAGTGGGTCAGACAGTCGAGTCG	CGGTAGTACGGGTTGGACGTTGGCCTTGTGA  A I M P N L Q T G T L  ACCTTCACCCAGTCTGTCAGCTCAGCGTACAC  TGGANGTGGGTCAGACAGTCGAGTCGCATGTG  T F T Q S V S S A Y T  GTGCCTATGATTATTGTCATCTTCTGCTACTT  CACGGATACTAATAACAGTAGAAGACGATGAA  V P M I I V I F C Y L  AGACGGAGGGTGAAACCCGACAACAAGCCCAA  AGACGGAGGGTGAAACCCGACAACAAGCCCAA  R R V K P D N K P K  GTCACCATGTTCGTAGTTTTTGTACTTTTTGCC  V T M F V V F V L F A  GGTCTTATTGTCGCCTCAGACCCTGCCACCAT  CCAGAATAACACCGGAGTCTGGGACCGTGGTA  G L I V A S D P A T M  GTGGCTAGTTACTACTACCTCCCGTACTTCAACAGC  841  GTGGCTAGTTACTACTACCTCCCGTACTTCAACAGCCAACAGCCCCAACAGCCCCACCAT  GTGGCTAGTTACTACCTCCCGTACTTCAACAGCCCTGCCACCAT  CCACCGATCAATAACTACCTCCCGTACTTCAACAGCCCTGCCACCAT  GTGGCTAGTTACTACCTCCCGTACTTCAACAGCCCTGCCACCAT  CACCGATCAATAACTACCTCCCGTACTTCAACAGCCCTGCCACCAT  GTGGCTAGTTACTACCTCCCGCTACTTCAACAGCCCTGCCACCAT  CACCGATCAATAATAGGACCCCCATGAAGTTGTCCCCCACCATCAACAGCCCCATCAACAGCCCTGCCACCATCAACAGCCCTGCCACCATCAACAGCCCTGCCACCATCAACAGCCCTGCCACCATCAACAGCCCTGCCACCATCAACAACAACAACAACAACAACACAGCCCTGCCACCATCAACAACAACAACAACAACAACAACAACAACAA	CGGTAGTACGGGTTGGACGTTTGGCCTTGTGAGGT  A I M P N L Q T G T L Q  ACCTTCACCCAGTCTGTCAGCTCAGCGTACACGAT  TGGAAGTGGGTCAGACAGTCGAGTCG	CGGTAGTACGGGTTGGACGTTTGGCCTTGTGAGGTCAT  A I M P N L Q T G T L Q Y  ACCTTCACCCAGTCTGTCAGCTCAGCGTACACGATAGC T F T Q S V S S A Y T I A  GTGCCTATGATTATTGTCATCTTCTGCTACTTAAGGAT  CACGGATACTAATAACAGTAGAAGACGATGAATTCCTA  V P M I I V I F C Y L R I  AGACGGAGGGTGAAACCCGACAACAAGCCCAAACTGAA  661 TCTGCCTCCCACTTTGCGCTGTTGTTCGGGTTTGACTT  R R R V K P D N K P K L K  GTCACCATGTTCGTAGTTTTTTGTACTTTTTGCCATTTG  721 CAGTGGTACAAGCATCAAAAACATGAAAAACGGTAAAC  V T M F V V F V L F A I C  661 CCAGAATAACACCGGAGTCTGGACCCTGCCACCATGGTCCC  781 CCAGAATAACACCGGAGTCTGGGACCGTGGTACCAGGGG  G L I V A S D P A T M V P  GTGGCTAGTTACTACCTGCCGTACTTCAACAGCTGCCT  841 CACCGATCAATGATGGACCGCATGAAGTTGTCGGACGGAACAACAAGCTCCCTTCAACAGCTGCCCTCCACCATGATCCCTCCC	CGGTAGTACGGGTTGGACGTTTGGCCTTGTGAGGTCATGCT  A I M P N L Q T G T L Q Y D  ACCTTCACCCAGTCTGTCAGCTCAGCGTACACGATAGCAGT  TGGAAGTGGGTCAGACAGTCGAGTCG	CGGTAGTACGGGTTGGACGTTTGGCCTTGTGAGGTCATGCTAGC  A I M P N L Q T G T L Q Y D P  ACCTTCACCCAGTCTGTCAGCTCAGCGTACACGATAGCAGTGGT  TGGAAGTGGGTCAGACAGTCGAGTCG	CGGTAGTACGGGTTGGACGTTTGGCCTTGTGAGGTCATGCTAGGGGG  A I M P N L Q T G T L Q Y D P R  ACCTTCACCCAGTCTGTCAGCTCAGCGTACACGATAGCAGTGGTGGT  TGGAAGTGGGTCAGACAGTCGAGTGCATGTGCTATCGTCACCACCA  T F T Q S V S S A Y T I A V V V V  GTGCCTATGATTATTGTCATCTTCTGCTACTTAAGGATATGGGTCCT  CACGGATACTAATAACAGTAGAAGACGATGAATTCCTATACCCAGGA  V P M I I V I F C Y L R I W V L  AGACGGAGGGTGAAACCCGACAACAAGCCCAAACTGAAGCCCCAGGA  AGACGGAGGGTGAAACCCGACAACAAGCCCAAACTGAAGCCCCAGGA  TCTGCCTCCCACTTTGGGCTGTTGTTCGGGTTTGACTTCGGGGTCCT  R R R V K P D N K P K L K P Q D  GTCACCATGTTCGTAGTTTTTGTACTTTTTGCCATTTGTTGGGCCCC  CAGTGGTACAAGCATCAAAAACATGAAAAAACGGTAAACAACCCGGGG  V T M F V V F V L F A I C W A P  GGTCTTATTGTGGCCTCAGACCCTGCCACCATGGTCCCCAGGATCCC  G L I V A S D P A T M V P R I P  GTGGCTAGTTACTACCTGGCGTACTTCAACAGCTGCCTCAACGCAAT  S41  GTGGCTAGTTACTACCTGGCGTACTTCAACAGCTGCTCCAACGCAAT  CACCCGATCAATGATGGACCGGATTGCTCCAACGCAATTACCTCGGGGTTGCGTTACTAACAGCTGCTCCAACGCAATTACCTGGCGTACTTCAACAGCTGCCTCAACGCAATTACCTGGCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGTACTTCAACAGCTGCCTCAACGCAATTACCTGCCGATGAAGTTGTCGACGGAGTTGCGTTACCACGCAATTACCTGCCGATGATGAAGAGTTGCCGTCAACGCAATTACCTGCCGATGATGAAGAGTTGCCGTCAACGCAATTACCCTGCCGATGAAGTTGTCGACGGAGTTGCGTTAACCAGCGAATTACCCCGAATGAAT	CGGTAGTACGGGTTGGACGITTGGCCTTGTGAGGTCATGCTAGGGGCCT  A I M P N L Q T G T L Q Y D P R I  ACCTTCACCCAGICTGTCAGCICAGCGTACACGATAGCAGTGGIGGTTT  TGGAAGTGGGTCAGACAGTCGAGTCGCATGTGCTATCGTCACCACCAAA  T F T Q S V S S A Y T I A V V V F  GTGCCTATGATTATTGTCATCTTCTGCTACTTAAGGATATGGGTCCTGGT  CACGGATACTAATAACAGTAGAAGACGATGAATTCCTATACCCAGGACCA  V P M I I V I F C Y L R I W V L V  AGACGGAGGGTGAAACCCGACAACAAGCCCAAACTGAAGCCCCAGGACTT  TCTGCCTCCCACTTTGGGCTGTTGTTCGGGTTTGACTTCGGGGTCCTGAA  R R R V K P D N K P K L K P Q D F  GTCACCATGTTCGTAGTTTTTGTACTTTTTGCCATTTGTTGGGCCCCACT  CAGTGGTACAAGCATCAAAAACATGAAAAAACGGTAAACAACCCGGGGTCA  V T M F V V F V L F A I C W A P L  GGTCTTTATTGTGGCCTCAGACCCTGCCACCATGGTCCCCAGGATCCCAGG  G L I V A S D P A T M V P R I P E  GTGGCTAGTTACTACCTGGCGTACTTCAACAGCTGCCTCAACGCAATTAT  CACCGATCAATGATGACCCGCATGAAGTTGTCGACGGATTCCAACGCAATTAT  CACCGATCAATGATGGACCGCATGAAGTTGTCGACGGAGTTGCGTTAATATA	CGGTAGTACGGTTGGACGTTTGGCCTTGTGAGGTCATGCTAGGGGCCCTAGA  A I M P N L Q T G T L Q Y D P R I Y  ACCTTCACCCAGTCTGTCAGCTCAGCGTACACGATAGCAGTGGTGGTTTTCC  TGGAAGTGGGTCAGACAGTCGAGTCG	CGGTAGTACQGGTTGGACGITTGGCCTTGTGAGGTCATGCTAGGGGCCTAGATGAA A I M P N L Q T G T L Q Y D P R I Y S  ACCTTCACCCAGTCTGTCAGCTCAGCGTACACGATAGCAGTGGTGGTTTTCCATT TGGAAGTGGGTCAGACAGTCGAGTCG	A I M P N L Q T G T L Q Y D P R I Y S C  ACCTTCACCCAGICTGCAGCTCAGCGTACACGATAGCAGTGGGGCCTAGATGAGGACA  A I M P N L Q T G T L Q Y D P R I Y S C  ACCTTCACCCAGICTGCTCAGCCTCAGCGTACACGATAGCAGTGGGGGTTTTCCATTTCATC  TGGAAGTGGGTCAGACAGTCGAGTCG

ATGRICTTTGTGGGGGGTTCAATGAAGAACAGATAAATTAATTTAGCCCTCTCCA 961+++++++ 1020 TACAAGAAACACCTCTCAAGTTTACTTCTTCGTCTATTCTAATTTACATTCGGGAGAGGT	
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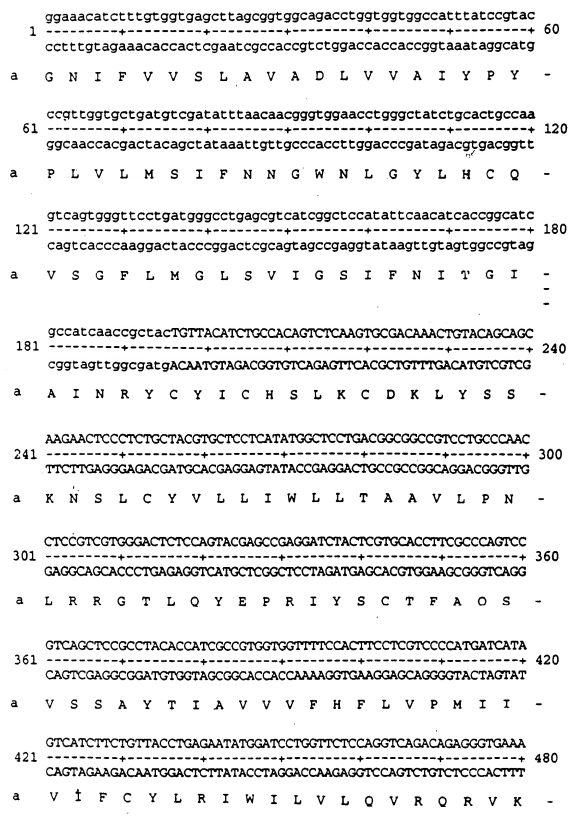


FIG. 4a

481	CC GG	rgac acty	CCG(	CAA GTT	ACC TGG	CAA GTT	ACT + TGA	GAA  CTT	ACC TGG	ACA TGT	CGA + GCT	CTI GAA	CAG GTC	GAA -+- CTT	TTT  AAA	TGT ACA	CAC + GTG	CAT GTA	GTI CĄĄ	TGTG + ACAC	540
a	P	D	R	K	P	ĸ	L	K	P	Н	D	F	R	N	F	V	T	M	F	V	-
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	AG	ACTY	GGG	GCG	GTC	GTA	CCA	.CGG	ATC	CTA	GGG	TCT	CAC	CGA	CAA	ACA	CCG	GTC	AAT	GATG	
a	S	D	P	A	S	М	V	P	R	I	P	E	W	L	F	V	A	S	Y	Y	-
661		GGC	GTA	TTT -+-	CAA				CAA									CCA	AAA′	TTTC	720
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	TC																				
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a		K																			-
	R AG	K	E TAA	Y CGA	r cgr	R GGC	I CGA	I TAG	V GGT	S TAA	L ATG	V GAA	T ACC	A GTC	R ICC	V ACTY	F	F GAC	V CAA	D CAAT	- 840
a 781	R AG	K	E TAA	Y CGA	R CGT	R	I CGA	I	V GGT	S TAA	L ATG	V GAA	T ACC	A GTC'	R	V ACTY	F GATY	F	V	D	- 840
	R AG	K	E TAA	Y CGA -+- GCT	R CGT	R GGC(	I CGA + GCT	I TAG  ATC	V GGT	S TAA  ATT	L ATG + TAC	V GAA  CTT	T ACC TGG	A GTC'	R ICC. AGG	V ACTY I'GA(	F GATY +	F GACO	V CAA( GTT(	D CAAT + GTTA	- 840 -
781 a	R AG TC S	K CTC GAG	E TAA  ATT N	Y CGA -+- GCT D	R CGT GCA V	R GGC( CCG( A	I CGA + GCT D	I TAG ATC R	V GGT CCA V	S TAA ATT K	L ATG + TAC W	V GAA CTT K	T ACC TGG	A GTC' -+- CAG	R ICC. AGG	V ACTY I'GA(	F GATY +	F GACO	V CAA( GTT(	D CAAT + GTTA	- 840 -
781	R AG TC S	K CTC GAG	E TAA ATT N	Y CGA GCT D	R CGT GCA V	R GGCG CCGG	I CGA GCT D	TAG ATC R	V GGT CCA V	TAA TAA ATT K	L ATG + TAC W	V GAA CTT K	T ACC TGG	A GTC' -+- CAG	R ICC. AGG	V ACTY I'GA(	F GATY +	F GACO	V CAA( GTT(	D CAAT + GTTA	- 840 -

FIG. 4b

	ATG	GCC	CTGC	:GGC	:CGG	GAC	GCG	IAAC	AGC	GA(	CCA'	ICC	AGG	GCA	ACG	ÇC	\GC(	3CG	CTG	CCC	'A
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С	v	v	v	F	н	F	L	v	P	М	I	I	v.	I	F	С	Y	L	R	I	-
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121	GG	GAC	CCA	-+- .CCG	AGC	STC	GCC	AC	AGG	CG	CCA	CGA	GTA	GCA	GT	GT	GGC	GGC	ACC	TGC	AGCAC	100
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		CAA	CT	CCT	GTC	GAT	CCT	CTC	CGI	GC?	CAC	GAZ	.ccc	CAA	GC1	cco	GAZ	ACGO	AGC	TAA'	r + 240	
181		GTT	GGA(	GGA(	CAC	CTA	GGA	GAG	GCA	CG2	AGTO	CII	GGC	GII	CG	\GG(	CTI	rgco	TCC	ATT		
a	G	N	L	L	v	I	L	S	V	L	R	N	R	K	L	R	N	A	G	N	-	
241		GTT	CTT	GGT	GAGT	rct	GGC	ATT	GGC	TG	ACCI	GGI	GGI	GGC	CTI	CTA	ACCC	CTA	.ccc	GCT/	A + 300	
241		CAA	GAA	CCA	CTC	AGA	CCG	TAA	CCC	AC:	rGGZ	CCA	CCA	CCG	GA?	GAI	GGG	GAT	GGG	CGA!		
a	L	F:	L	v	s	L	A	L	A	D	L	V	V	A	F	Y	P	Y	P	L	-	
301		CCT	CGT	GGC(	CATO	TT	CTA	TGA	CGG	CTC	GGC	CCT	GGG	GGA	.GGA	GCA	CTG	CAA	GGC	CAGO	360	
	TA	GGA(	GCA(	CCG	GTAC	GAA	GAT.	ACT	GCC	GAC	ccc	GGA	.CCC	CCT	CCT	CGI	'GAC	GTT	CCG	GTC	3	
a	I	L	V	A	I	F	Y	D	G	W	A	L	G	Ε	Ε	Н	С	K	A	S	-	
361		CTT	TGT	GAT		CCT		CGT			GCT(				TAT	CAC	TGC			CAT	r + 420	
301		GAA	ACA	CTA	CCC	GGA	CTC	GCA	GT?	\GC(	CGAC	SACA	\GA.	GTT	ATA	GTC	ACC	GTA	GCG	GTA	A	
a	A	F	V	М	G	L	S	V	I	G	S	V	F	N	I	T	A	I	A	I		
421				-+-			+				- +			-+-			+				+ 480	
_																				CGT		,
a	N	Ŕ	Y	С	Y	I	С	Н	S.	M	A	Y	Н	R	I	Y	R	R	W	H.	-	

FIG. 6a

4.	481				-+-			+				+			-+-						CTTC + GAAG	540
	a	Т	P	L	Н	I	С	L	I	W	L	L	T	V	V	A	L	L	P	N	F	-
	541		TGT	GGG	GTC	CCI	GGA	GTA	CG.	ACCO	CACC	CAI	CTA	ATTO	CTC	CAC	CT1	CAT	rccz	GAC	CGCC	600
	2																				GCGG	
,	a	r	V	G	S	L	E	Y	ט	Þ	R	1	Y	5	C	T	F	I	Q	Т	A	-
	6 <b>01</b>				-+-			+				+			-+-			+				660
		TC	GTG	GGT	CAT	GTG	CCG	CCG	TCA	CCA	.CCA	GTA	GGT	'GAA	GGA	.GGA	GGG	АТА	.GCG	ACA	GCAC	
	a	S	T	Q	Y	T	A	A	V	V	V	I	Н	F	L	L	P	I	A	V	V	
	6 <b>6</b> 1				-+-			+				+			-+-			+			•	720
	a		gaa F	C							L L										'CGGT P	-
	721		'-		+-			+				+			+-						GGTG + CCAC	780
	a	E	s	R	L	С	L	K	P	S	D	L	R	S	F	L	Т	М	F	V	V	-
	7 <b>81</b>				-+-			+				+			-+-			+			CATC + GTAG	840
	a	F	v	I	F	A	I	С	W	A	P	L	N	С	I	G	Ļ	Α	v	A	I	-
	8 <b>4</b> 1				-+-			+				+			-+-			+			ACTG + TGAC	900
	a	N	P	Q	E	М	A	P	Q	I	P	E	G	L	F	v	Т	s	Y	L	L	-

901	GCTTATTTCAACAGCTGCCTGAATGCCATTGTCTATGGGCTCTTGAACCAAAACTTCCGC	ACA(	SCT.	GCCI	rgaa 	TGC	CAT	TGT	CTA	lgg.	gCT	CHI	SAAC	CAZ	AAAC	)TT	2000	050
	CGAATAAAGTTGTCGACGGACTTACGGTAACAGATAcccGAGAACTTGGTTTTGAAGGCG	TGT	CGA	තිවුට	ACTT	ACG	GTA	ACA	GAT	ACC	SG.	GAA	) TT	GT.	Ĭ	3AA(	3366	200
ď	AYFNSCLNAIVYGLLNQNF	Ŋ	U	i i	Z	æ	н	>	>-	O	Ţ	L	z	ø	z	្រុ	ፎ	1
061	-	AGAG	3GA.	ICCI	CIT	ည	CCT	TTG	3AAC	CCC	ACG.	3CAC	TGC.	ATT	Ç	\GA7	GCT	
700	TCCCTTATGTTCTCCTAGGAGAACCGGGAAACCTTGGGTGCCGTGACGTAAGTTCTACGA	Si Si	i i	AGGA	GAA	500	GGA	AAC(		55	- <u>2</u> 2	GTC	ACG	TAA	GTT	CLZ	++	1020
ಹ	REYKRILLALWNPRHCIQDA	弘	H	IJ	u	4	IJ	3	z	Ω,	r.	I	O	н	ø	Ω	Æ	ı
1021	TCCAAGGGCAGCCACGCGGAGGGCTTGCAGAGCCCAGCTCCACCCATCATTGGTGTGCAG 	2	9 0	4500 4500	999	GCT	SCA G	SAG TE		1001 1001 1001	δ + 6		ATC 	ATT -+-	GGT	GTG	CAG +	1080
ಹ	SKGSHAEGLOS	H	A A	) ш	) (5)	<u>,</u>	3 0	S	P A P P I I G V	<b>4</b>	<u>д</u>		Н	Н	5 0		<b>j</b> 0	
1001		\TGC	ĬŢ.	ICTA	သည	JG	(6	EO	(SEQ ID NO:15)	õ	[2]							
1001	GTGGTCCGTCTACGAGAGATCGGAC	rACG	AG.	AGAT	2000	AC	011	n										
ಹ	H Q A D	D A L	J	*		(SE	0	0	(SEQ ID NO:16)	_								

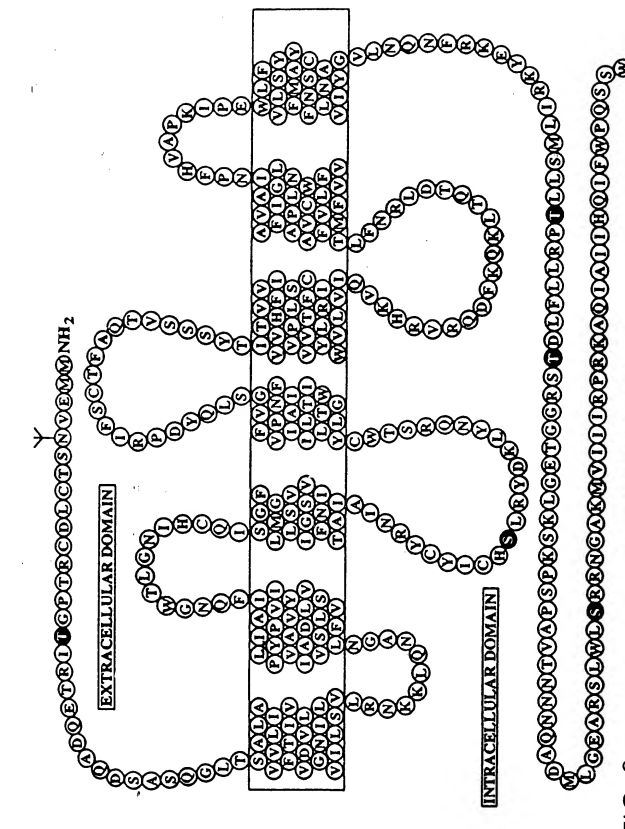
FIG. 6c

xmr	MMEVNSTCLDCRTPGTIRTEQDAQDSASQGLT MAGRLWGSPGGTPKGNGSSALLNVSQAAPGAGDGVRPRPSWLA I
xmr	SALAVVLIFTIVVDVLGNILVILSVLRNKKLQNAGNLFVVSLSIADLVVAVYPYPVILI
ov	ATLASILIFTIVVDIVGNLLVVLSVYRNKKLRNAGNVFVVSLAVADLLVAVYPYPLALA
hum	GNIFVVSLAVADLVVAIYPYPLVLM
xmr	AIFONGWTLGNIHCOISGFLMGLSVIGSVFNITAIAINRYCYICHSLRYDKLYNQRSTW
ov	SIVNNGWSLSSLHCOLSGFLMGLSVIGSVFSITGIAINRYCCICHSLRYGKLYSGTNSL
hum	SIFNNGWNLGYLHCQVSGFLMGLSVIGSIFNITGIAINRYCYICHSLKCDKLYSSKNSL
xmr	CYLGLTWILTIIAIVPNFFVGSLQYDPRIFSCTFAQTVSSSYTHTVVVVHFIVPLSVVT
ov	CYVFLIWTLTLVAIVPNLCVGTLQYDPRIYSCTFTQSVSSAYTHAVVVFHFIVPMLVVV
hum	CYVLLIWLLTA. AVLPNLRRGTLQYEPRIYSCTFAQSVSSAYTHAVVVFHFLVPMIIVI
xmr	ECYLRIWVLVIQVKHRVRQDFKQKLTQTDLRNFLTMFVVFVLFAVCWAPLNFIGLAVAI
ov	FCYLRIWALVLQVRWKVKPDNKPKLKPQDFRNFVTMFVVFVLFAICWAPLNFIGLVVAS
hum	FCYLRIWILVLQVRQRVKPDRKPKLKPHDFRNFVTMFVVFVLFAICWAPLNFIGLAVAS
xmr	NPFHVAPKI PEMLFVLSYFMAYFNSCLNAVIYGVLNQNFRKEYKRILMSLLTPRLLFLD
ov	DPASMAPRI PEMLFVASYYMAYFNSCLNAI IYGLLNQNFRQEYRKI IVSLCTTKMFFVD
hum	DPASMVPRI PEMLFVASYYMAYFNSCLNAI ISGYWNQNFRKEYRRI IVSLVTARVFFVD
xmr	TSRGGTEGLKSKPSPAVTNNNQADMLGEARSLWLSRRNGAKMVIIIRPRKAQIAIIHQIF
ov	SSNHVADRIKRKPSPLIANHNLIKVDSV* SEQ ID NO:4
hum	SSNDVARDVKWKPSPLMTNNNVVKVDSV* SEQ ID NO:6
xmx	WPQSSWATCRODTKITGEEDGCRELCKDGISQR SEQ ID NO:2 FIG. 7

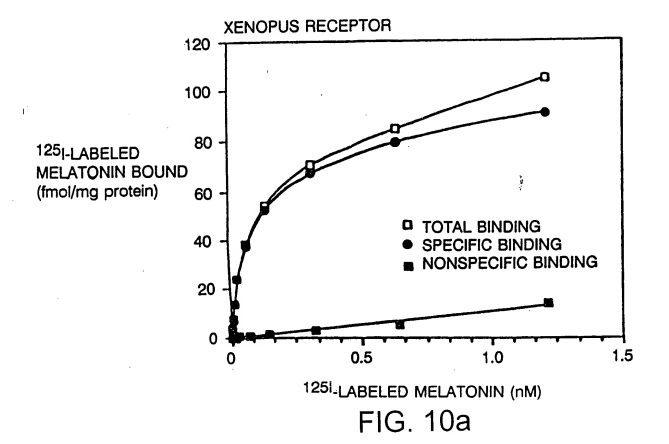
QIFWPQSSWATCRQDTKITGEEDGCRELCKDGISQR 420 (SEQ ID NO:2)

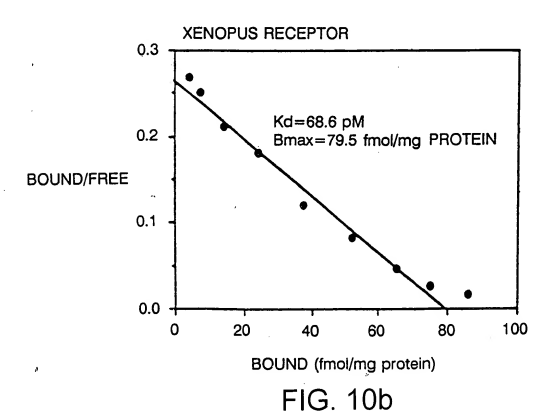
Xenopus

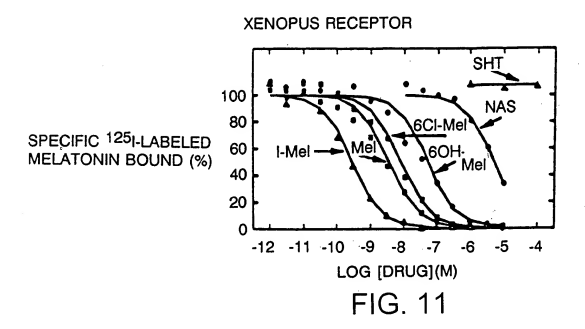
VVLSVYRNKKLRNAGN 79 VILSVYRNKKLRNAGN 63 VILSVLRNKKLQNAGN 68	/-LSV-RNKKL-NAGN	CCICHSLRYGKLYSGT 158 SYICHSLKYDKLYSSK 142 SYICHSLRYDKLYNQR 147	ICHSL-Y-KLY	YLRIWALVIQVRWKV 237 YYLRIWILVIQVRQRV 221 YYLRIWVLVIQVKHRV 226	YLRIW-LV-QVV	I FNSCLNAIIYGLLNQ 316 FNSCLNAIIYGLLNQ 300 FNSCLNAVIYGVLNQ 305	FNSCLNA-IYG-LNQ	(SEQ ID NO:4) (SEQ ID NO:12) RRNGAKHVIIIRPRKAQIAIH 334
I MAGRLWGSPGGTPKGNGSSALLNYSQAAPGAGDGVRPRPSWLAATLASILIFTIVVDIVGNLLVVLSVYRNKKLRNAGN MQGNGSALPNASQPVLRGDGARPSWLASALACVLIFTIVVDILGNLLVILSVYRNKKLRNAGN MMEVNSICLDCRTPGTIRTEQDAQDSASQGLTSALAVVLIFTIVVDVLGNILVILSVLRNKKLQNAGN		II VFVVSLAVADLLVAVY PY PLALASIVNNGWSLSSLHCQLSGFLMGLSVIGSVFSITGIAINRYCCICHSLRYGKLYSGT IFVVSLAVADLVVAIY PY PLVLMSIFNNGWNLGYLHCQVSGFLMGLSVIGSIFNITGIAINRYCYICHSLKYDKLYSSK LFVVSLSIAADLVVAVY PY PVILIAIFQNGWTLGNIHCQISGFLMGLSVIGSVFNITAIAINRYCYICHSLRYDKLYNQR	-FVVSLADL-VA-YPYPLINGW-LHCQ-SGFLMGLSVIGS-F-IT-IAINRYC-ICHSL-Y-KLY	NSLCYVFL IWTLTLVA IV PNLCVGTLQYDPR IY SCTFTQ SVSSAYT IAVVVFHF IV PMLVVVFCYLR IWALVLQVRWKV NSLCYVLL IWLLTLAAVL PNLRAGTLQYDPR IY SCTFAQ SVSSAYT IAVVVFHFLVPM I IV IFCYLR IWILVLQVRQRV STWCYLGLTWILTIIAIV PNFFVGSLQYDPR IFSCTFAQTVSSSYT ITVVVVHFIV PLSVVTFCYLR IWVLVIQVKHRV	ConsensusCYL-W-LTAPNG-LQYDPRI-SCTF-Q-VSS-YTI-VVV-HF-VPV-FCYLRIW-LV-QVV	VII KPDNKPKLKPQDFRNFVTMFVVFVLFAICMAPLNFIGLVVASDPASMAPRIPEWLFVASYYMAYFNSCLNAIIYGLLNQ KPDRKPKLKPQDFRNFVTMFVVFVLFAICWAPLNFIGLAVASDPASMVPRIPEWLFVASYYMAYFNSCLNAIIYGLLNQ RQDFKQKLTQTDLRNFLTMFVVFVLFAVCWAPLNFIGLAVAINPFHVAPKIPEWLFVLSYFMAYFNSCLNAVIYGVLNQ	D-K-KL-P-D-RNF-TMFVVFVLFA-CWAPLNFIGL-VAPP-IPEWLFV-SY-MAYFNSCLNA-IYG-LNQ	TKMFEVDSSNHVADRIKRKPSPLIANHNLIKVDSV 366 ARVFEVDSSNDVADRVKWKPSPLMTNNNVVKVDSV 350 PRLLFLDTSRGGTEGLKSKPSPAVTNNNQADMLGEARSLWLSI
Sheep M Human Xenopus	Consensus	Sheep vi Human Ii Xenopus Li	Consensus -	Sheep NS Human NS Xenopus ST	Consensus -	Sheep Ki Human Ki Xenopus R	Consensus -	Sheep NFRQEYRKIIVSLCTI Human NFRKEYRRIIVSLCTI Xenopus NFRKEYKRILMSLLTI

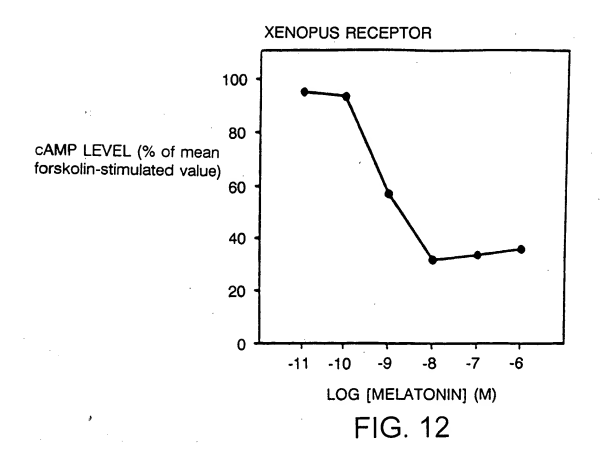


COOH (ROST DED KICLE BICCE E CETOKED DE BEST COOH (ROST DE BEST COOH ( FIG. 9 seq id no:2



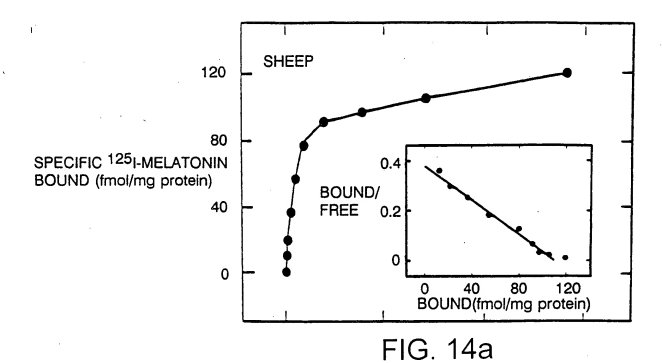


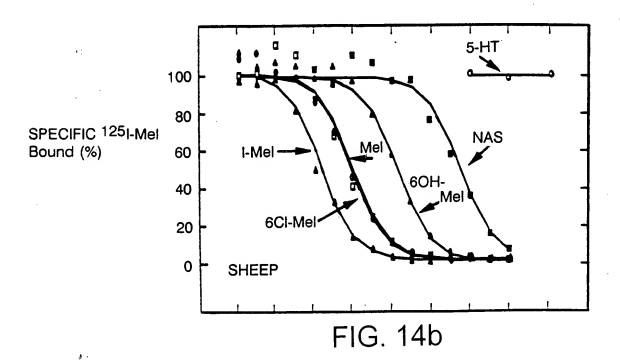




# Kb 7.5 — 4.4 — 2.4 —

1.4 — FIG. 13





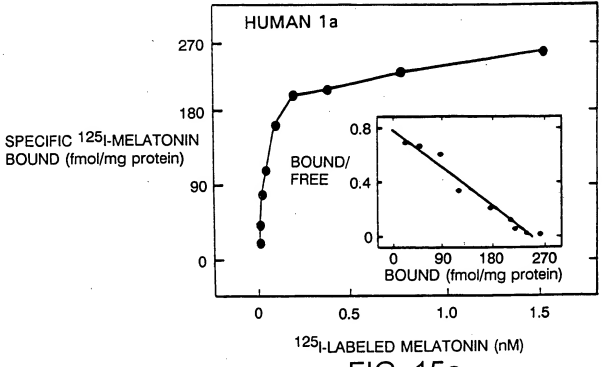


FIG. 15a

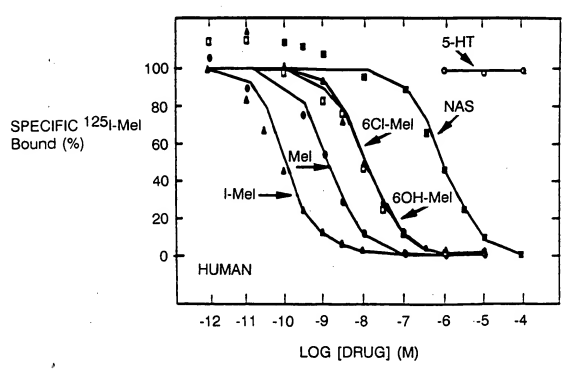


FIG. 15b

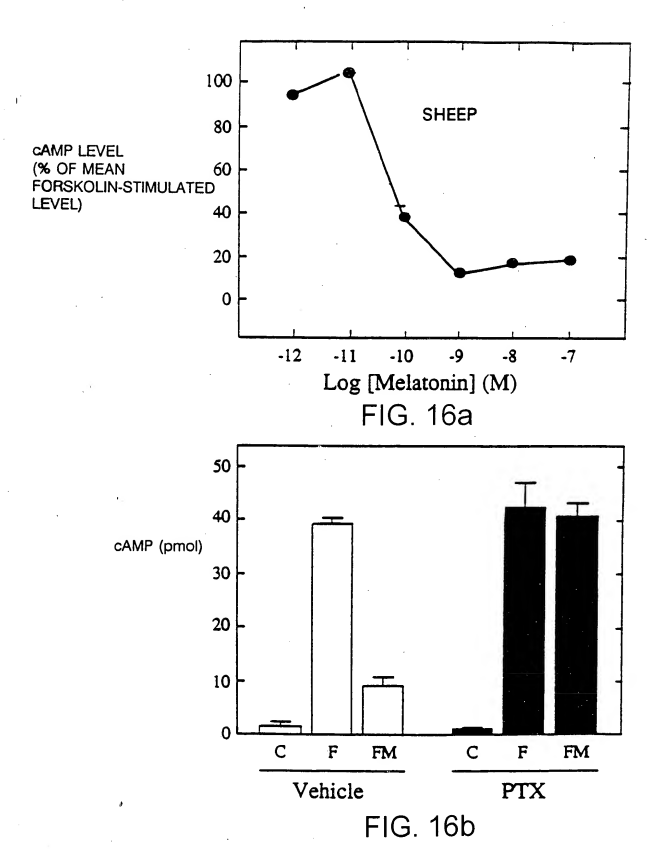


FIG. 17a

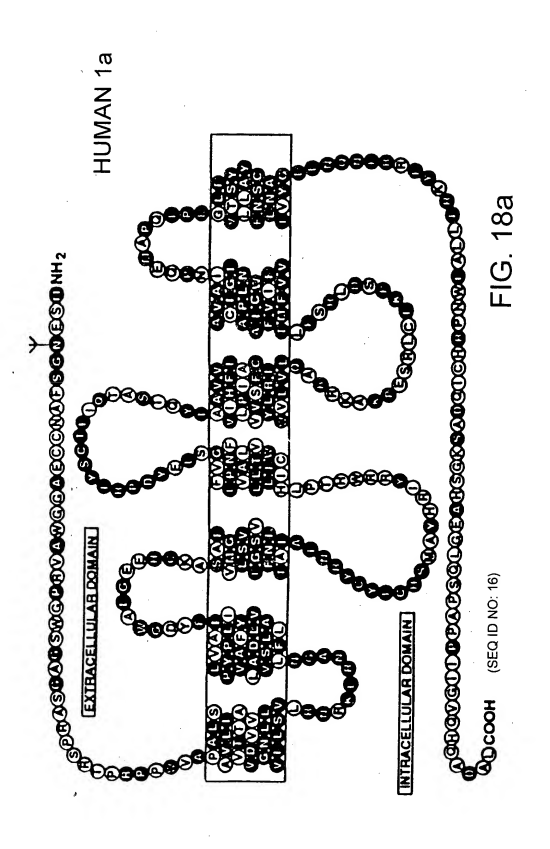
FIG. 17b

125I-MEL Binding

Histology

mRNA

FIG. 17c



Human 1b  HEVASICECEAGGANAPPCHSGAGSARPSRTPRPPMADALSAVLIVITAVDVOGNILVILSVIRNKIRNAGNIEVSLALADLIVATFYPTPLILVAIFYDCHAL  Kanopus  HEVASICLDCRITPCTIRTEQDAQDSASQCLISALAVVLIFTTVVDICANLIVILSVRNKKIRNAGNIEVSLALADLIVATYPYPTVLILATIFQRGNTL  Consensus  LII  CEHCKASAFVACLSVIGSTENTTALAINNYCYICHSHAYRNTRRHITPLAICLINLITAVALPHFFVGSLEYDPRISCTFAQTVSSTATITAVVVHFLIPINAVSFC  Kanopus  G-HC-S-F-HGLSVIGSTENTTALAINNYCYICHSLAYDRINTRRHITPLAICLINLITALAVLPHFFVGSLEYDPRISCTFAQTVSSSTATITAVVVHFLIPINAVFFC  CONSENSUS  G-HC-S-F-HGLSVIGSTENTTALAINNYCYICHSLAYDRINTRRHITPLAICLINLITALAVLPHTATSTATVAVHFLIPINAVFFC  CONSENSUS  TALHIHULVIQARRAKAPESRICIKPSDLRSFLTHFVVFVIFAIGNAPLHFTGLAVAINTPRETRYSTILAATHYSCI	Human 1b Human 1a Xenopus Consensus Human 1a Xenopus Consensus Y Kuman 1a Xenopus Consensus Y Consensus Y Consensus Y Consensus Y Consensus Y Consensus Y
ConsensusD-S	Consensus -
IQDASKGSHAEGLQSPAPPIIGVQHQADAL 362 (SEQ 1D NO:18) TVDSSNDVADRVKWKPSPLYTNNNVVKVDSV 350 (SEQ 1D NO:12) TLDTSRGGTEGLKSKPSPAVTNNNQADMLGEARSLWLSRRNGAKHVIIIRPRKAQIAIIHQIFWPQSSWATCRQDTKITGEEDGCRELCKDGISQR 420	78
ltw-lv-q	Consensus Y
% INTILVIQYRRYKPDRKPKLKPQDFRNFVIFY VFV IF ALLWAFLNF IGLAVASIPPEGIFVTSYLLAYFNSCLNAIVYGLLNONFRREYKFILLALWNPR NIWLVIQVKHRVRQDFKQKLTQTDLRNFLTHFVVFVLFAICWAPLNF IGLAVASDPASHVPRIPEWLFVASYYHAXFNSCLNAIIYGLLNQNFRKEYRRIIVSLCTAR NIWLVIQVKHRVRQDFKQKLTQTDLRNFLTHFVVFVLFAVCWAPLNF IGLAVAINPFHVAPKIPEWLFVLSYFHAYFNSCLNAVIYGVLNQNFRKEYKRILHSLLTPR	Human la Y Xenopus Y
VII NIWLVLQARRKAKPESRLCLKPSDLRSFLTHFVVFVIFAICHAPLNCIGLAVAINPOFMABOIDFGI FUTEVI I AVENECI IN TUVOTI	-
-HCS-F-MGLSVIGS-FNIT-IAINRYCYICHSYL-W-LTAPNG-L-YDPRI-SCTF-QSYTVVV-HFPV-FC	Consensus G
LHCQVSGFLMGLSVIGSVFNITALAINNYCYICHSMAYHHIYRWHTPLHICLIMLLTVVALLPNFFVGSLEYDPRIYSCTFIQTASTQYTAAVVVIHFLLPIAVVSFC LHCQVSGFLMGLSVIGSIFNITGIAINNYCYICHSLKCDKLYSSKHSLCYVLLIMLLTLAAVLPNLRAGTLQYDPRIYSCTFAQSVSSAYTIAVVVFHFLVPHIIVIFC IHCQISGFLMGLSVIGSVFNITAIAINNYCYICHSLRYDKLYNQRSTWCYLGLTWILTIIAIVPNFFVGSLQYDPRIFSCTFAQTVSSSYTITVVVVHFIVPLSVVTFC	Human la C Xenopus G
	Consensus
nserlijs anchemasmavrrengsagsarpsktprppwapalsavlivttavdvvgnillvilsvirnkurnagnifloslaladilvafypyliilvaifydchal MQG <u>NGS</u> ALP <u>NASQ</u> PVLRGDGARPSWLASALACVLIFTTVVDILGNILVILSVYRNKKLRNAGNIFVVSLAVADLVVAIYPYPLVILASIFNNGWNL MMEVNSICLDCRTPGTIRTEQDAQDSASQGLISALAVVLIFTTVVDVLGNILVILSVLRNKKLQNAGNIFVVSLSIADLVVAVYPYPVILIAIFQNGWTL	Human la Xenopus
II	Human 1b

FIG. 18b

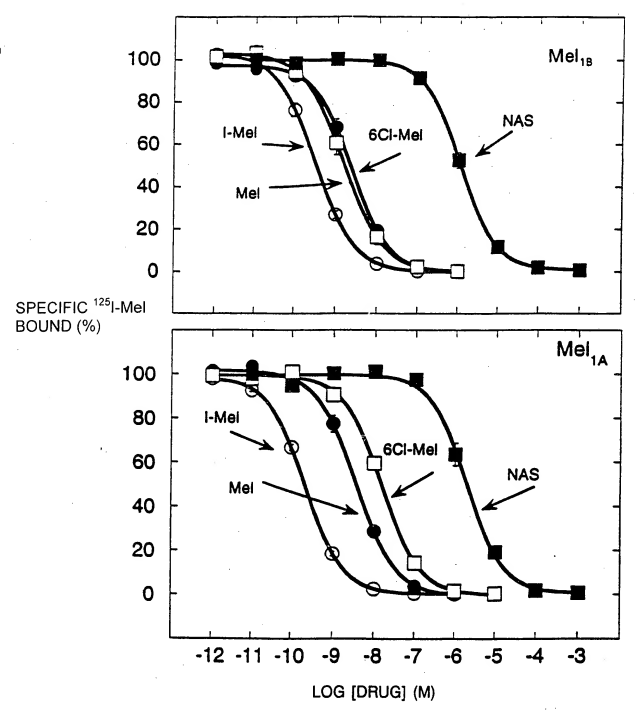
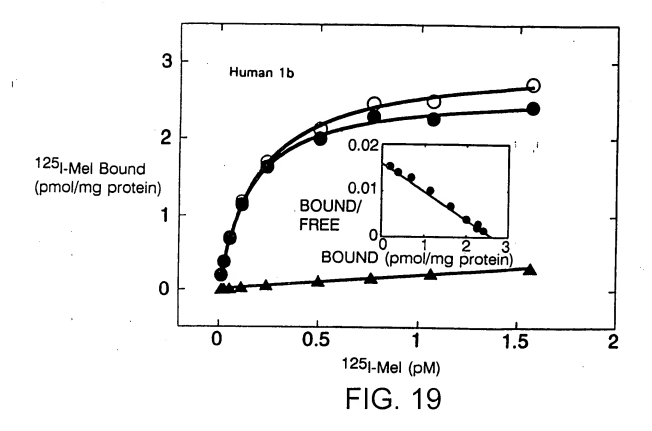
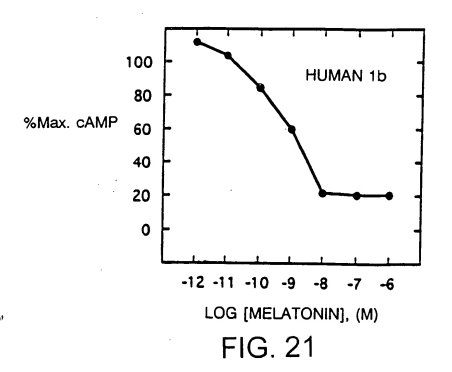


FIG. 20







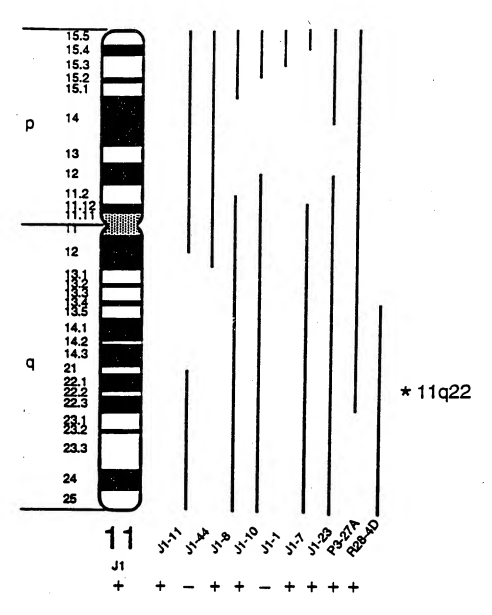


FIG. 23

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